

**PATIENT LEAFLET IN ACCORDANCE  
WITH THE PHARMACISTS'  
REGULATIONS (PREPARATIONS) – 1986**

This medicine is dispensed without a  
doctor's prescription

**Tiptipot Simicol**



**Oral drops**

The active ingredient and its concentration:  
Simethicone 20 mg/0.3 ml (66.67 mg/1 ml)  
Inactive ingredients and allergens in the  
preparation – see the subsection "Important  
information about some of the ingredients of the  
medicine" and section 6.

**Read the entire leaflet carefully before using  
the medicine.** This leaflet contains concise  
information about the medicine. If you have  
any other questions, refer to the doctor or the  
pharmacist.

Use the preparation according to the  
instructions in the dosage section of this leaflet.  
Consult the pharmacist if you need further  
information. Refer to the doctor if signs of the  
ailment (symptoms) worsen or do not improve.

**1. What is the medicine intended for?**

The medicine is intended for the relief of  
abdominal pain in infants and young children  
caused by the accumulation of gas.

**Therapeutic class:** medicines for the relief of  
gastrointestinal gas.

**2. Before using the medicine:**

**✗ Do not use this medicine if:**

- There is a known sensitivity (allergy) to  
the active ingredient or to any of the other  
ingredients this medicine contains (see  
section 6).

**✗ Drug interactions:**

**If the child is taking or has recently taken  
other medicines, including non-prescription  
medicines and nutritional supplements, tell  
the doctor or pharmacist.** Especially if the  
child is taking:

- Levothyroxine for treatment of thyroid  
disorders. Tiptipot Simicol may reduce the  
amount of absorbed levothyroxine and  
diminish its effect.

**✗ Pregnancy, breastfeeding and fertility:**

Tiptipot Simicol is intended especially for  
children. If you are an adult woman using the  
medicine, consult a doctor or pharmacist before  
starting treatment if you are pregnant, may be  
pregnant, are planning to become pregnant or  
are breastfeeding.

Do not use the preparation during pregnancy  
unless the doctor tells you to.

**✗ Important information about some of the  
ingredients of the medicine:**

The medicine contains 1.2 mg of sodium  
benzoate per 0.6 ml. Sodium benzoate may  
increase jaundice (yellowing of the skin and  
eyes) in infants up to 4 weeks of age. The  
medicine contains less than 23 mg of sodium  
per dose and is therefore considered sodium-free.

**3. How should you use the medicine?**

Check with the doctor or pharmacist if you are  
uncertain about the dosage and how to use the  
preparation.

The generally accepted dosage is:

0.3 ml (20 mg) before each meal. The  
dosage can be increased to 0.6 ml (40 mg) if  
necessary.

Do not exceed 12 doses of 0.3 ml or 6 doses of  
0.6 ml (240 mg in total) per day.

During the period of treatment with the  
medicine, it is advisable to consult a doctor for  
the purpose of diagnosing the cause of gas  
accumulation.

**Do not exceed the recommended dose.**

**Method of use:**

1. Shake well before use.
2. Fill the enclosed syringe to the desired  
amount and drip into the mouth.



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3. After use, separate the parts of the syringe  
and rinse the two parts of the syringe  
thoroughly with lukewarm water.

With liquid medicines, use the syringe or  
dropper intended for measuring the correct  
amount of medicine. If a measuring device  
is not included in the package, consult a  
pharmacist. Do not use a household teaspoon  
to measure the amount of medicine. Household  
teaspoons vary in size and it is likely you will  
not receive the correct amount of medicine.

**If the child has taken an overdose** or if a  
child has accidentally swallowed the medicine,  
proceed immediately to a doctor or a hospital  
emergency room and bring the package of the  
medicine with you.

**If you forget to give the medicine** at the  
required time, do not give a double dose.  
Administer the next dose at the regular time  
and consult a doctor.

**Do not take medicines in the dark! Check  
the label and the dose every time you take a  
medicine. Wear glasses if you need them.  
If you have any other questions regarding  
the use of the medicine, consult the doctor  
or the pharmacist.**

**4. Side effects:**

As with any medicine, using Tiptipot Simicol  
may cause side effects in some users. Do not  
be alarmed when reading the list of side effects.  
The child may not suffer from any of them.  
Possible side effects are nausea, constipation  
or an allergic reaction such as skin rash, itching  
of the skin, swelling of the face or tongue or  
difficulty breathing.

**If a side effect occurs, if one of the side  
effects worsens, or if you suffer from a side  
effect not mentioned in this leaflet, consult  
your doctor.**

**Reporting side effects:**

Side effects may be reported to the Ministry  
of Health by clicking on the link "Report side  
effects due to medicinal treatment" found on the  
Ministry of Health website homepage  
([www.health.go.kr](http://www.health.go.kr)), which will direct you to  
the online form for reporting side effects, or by  
clicking on the following link:  
<https://sideeffects.health.go.kr/>

**5. How to store the medicine?**

Avoid poisoning! This medicine and any other  
medicine must be kept in a closed place out of  
the reach and sight of children and/or infants to  
avoid poisoning. Do not induce vomiting without  
an explicit instruction from the doctor.

Do not use the medicine after the expiry date  
(exp.) appearing on the package. The expiry  
date refers to the last day of that month.  
Store at a temperature below 25°C.  
Once the bottle has been opened, Tiptipot  
Simicol may be used for 2 months.

**6. Additional information:**

In addition to the active ingredient, the medicine  
also contains:

Polysorbate 80, Sorbitan Monostearate 60,  
Microcrystalline cellulose, Strawberry Flavour,  
Citric acid (anhydrous), Sodium Benzoate,  
Sodium Citrate, Xanthan Gum, Saccharin  
sodium, Purified Water.

**What does the medicine look like and what  
are the contents of the package:**

A glass bottle containing 30 ml of a white and  
viscous strawberry flavored emulsion.

**Name of manufacturer/marketing  
authorization holder and address:** CTS  
Chemical Industries Ltd., 3 Hakidma st., Kiryat  
Malachi.

This leaflet was revised in 06/2022 in  
accordance with the Ministry of Health  
guidelines.

Registration number of the medicine in the  
national drug registry of the Ministry of Health:  
533826579

